5' GCC AGC TGC GTT CTG AGC CTG GGC GCA GCT ACC ATC TGC TCT GGG AAG CAC CAG GGT GTC CCC GCC GCC CTC AGC TCG AAG TCA GCC ACC ATG GAG GCG CAG GCA CAA MEAQAQ GGT TTG TTG GAG ACT GAA CCG TTG CAA GGA ACA GAC GAA GAT GCA GTA GCC AGT G L L E T E P L Q G T D E D A V A S GCT GAC TTC TCT AGC ATG CTC TCT GAG GAG GAA AAG GAA GAG TTA AAA GCA GAG F S S M L S E E E E E L K A E TTA GTT CAG CTA GAA GAC GAA ATT ACA ACA CTA CGA CAA GTT TTG TCA GCG AAA Q L E D E I T T L R Q V L S A K GAA AGG CAT CTA GTT GAG ATA AAA CAA AAA CTC GGC ATG AAC CTG ATG AAT GAA ERHLVEIKQKLGMNLMNE TTA AAA CAG AAC TTC AGC AAA AGC TGG CAT GAC ATG CAG ACT ACC ACT GCC TAC LKQNFSKSWHDMOTTTA AAG AAA ACA CAT GAA ACC CTG AGT CAC GCA GGG CAA AAG GCA ACT GCA GCT TTC K K T H E T L S H A G Q K A T A A AGC AAC GTT GGA ACG GCC ATC AGC AAG AAG TTC GGA GAC ATG AGT TAC TCC ATT SNVGTAIS K K F G D CGC CAT TCC ATA AGT ATG CCT GCT ATG AGG AAT TCT CCT ACT TTC AAA TCA TTT SISMPAMRNSPT F K S GAG GAG AGG GTT GAG ACA ACT GTC ACA AGC CTC AAG ACG AAA GTA GGC GGT ACG ERVETTVTSLKTKVGG AAC CCT AAT GGA GGC AGT TTT GAG GAG GTC CTC AGC TCC ACG GCC CAT GCC AGT NPNGGSFEEVLSSTAHA GCC CAG AGC TTG GCA GGA GGC TCC CGG CGG ACC AAG GAG GAG GAG CTG CAG TGC A Q S L A G G S R R T K E E E L O C

FIGURE 1A

767 776 785 NTG CCT GTG CTT ATC CAG ATA AGA AGA CCA AA 3'

FIGURE 1B

10. 5' TMG MKC GCG GGC CCC CGC CAG TCA GGT GGG TGC CAG GCC CTG GCC GTG GCG AAA GAG CCG GCG GAG GGA GGA CCC GCT CCC GGA GAC GCC GCC TCG CGA TCC CCG CGC GGG CGG GAC CGG GCC GGC ATC ATG ACC CTG TTT CAC TTC GGG AAC TGC TTC TLFHF GCT CTT GCC TAC TTC CCC TAC TTC ATC ACC TAC AAG TGC AGC GGC CTG TCC GAG ALAYFPYFITYKC TAC AAC GCC TTC TGG AAA TGC GTC CAG GCT GGA GTC ACC TAC CTC TTT GTC CAA A F W K C V QAGVTYL CTC TGC AAG ATG CTG TTC TTG GCC ACT TTC TTT CCC ACC TGG GAA GGC GGC ATC F L A T F F P T W E G G TAT GAC TTC ATT GGG GAG TTC ATG AAG GCC AGC GTG GAT GTG GCA GAC CTG ATA F I G E F M K A S V D V A D L GGT CTA AAC CTT GTC ATG TCC CGG AAT GCC GGC AAG GGA GAG TAC AAG ATC ATG N L V M S R N A G K G E Y K I GIT GCT GCC CTG GGC TGG GCC ACT GCT GAG CTT ATT ATG TCC CGC TGC ATT CCC V A A L G W A T A E L I M S R C I CTA TGG GTC GGA GCC CGG GGC ATT GAG TTT GAC TGG AAG TAC ATC CAG ATG AGC LWVGARGIEFDWKY ATA GAC TCC AAC ATC AGT CTG GTC CAT TAC ATC GTC GCG TCT GCT CAG GTC TGG S N I S L V H Y I V A S A Q ATG ATA ACA CGC TAT GAT CTG TAC CAC AAC TTC CGG CCA GCT GTC CTT CTG CTG TRYDLYHNF.RPAVLL ATG TTC CTC AGT GTC TAC AAG GCC TTT GTT ATG GAG ACC TTC GTC CAC CTC TGC LSVYKAFVMETFV TCG CTG GGC AGT TGG GCA RCT CTA MTG GCC CGA GCA GTG GTA ACG GGG CTG CTG G S W A X L X A R A V V T G L L

FIGURE 2A

793 802 GCC CTC AAG CAC TIT GGS CCT GTA TGT CGS CGT TGT CAA TGT GCA CTY CTA GGC A L K H F G P V C R R C Q C A L L G TTG GTG TCT CAG ACA TTG ATG TAC CTT TTC CCT GCC TCA CTC CAG GTT TTA GTG L V S Q T L M Y L F P A S L Q V L V AAG TAA ACA GTA TTT GGA AAG TT 3'

Library	Lib Description	Abun	Pct Abun
BRAINOM02	brain, 55 M, NORM, WM	1	0.0454
UTRSNOT01	uterus, 59 F	1	0.0393
TLYMNOR01	lymphocytes (non-adher PBMNC), 24 M, RP	1	0.0372
BRSTNOT02	breast, 55 F, match to BRSTTUT01	2	0.0317
PROSTUT08	prostate tumor, 60 M, match to PROSNOT14	1	0.0266
PROSNOT14	prostate, 60 M, match to PROSTUTO8	1	0.0256
LIVRNOM01	liver, 49 M, WM	1	0.0254
PROSNOT15	prostate, 66 M, match to PROSTUT10	1	0.0241
NERVMSM01	multiple sclerosis, 46 M, NORM, WM	1	0.0228
HNT2AGT01	hNT2 cell line, post-mitotic neurons	1	0.0190
BRAITUT02	brain tumor, metastasis, 58 M	1	0.0169
LIVSFEM02	liver/spleen, fetal M, NORM, WM	2	0.0053

Electronic Northern Results returned a total of 12 row(s).

FIGURE 3

Library	Lib Description	Abun	Pct Abun
COLNNOT22	colon, 56 F		
COLNPOT01			2 0.0554
PROSNOT18	prostate, hyperplasia, 58 M		2 0.0513
MUSCNOT02	muscle, psoas, 12 M		2 0.0513
STOMTUT01	stomach tumor, 52 M, match to STOMNOTO2		1 0.0382
SINTNOT02	small intestine, 55 F		1 0.0368
LVENNOT03	heart, left ventricle, 31 M		1 0.0345
MMLR3DT01	macrophages (adher PBMNC), M/F, 72-hr MLR		1 0.0337
PROSTUT01	prostate tumor, 50 M, match to PROSNOTO2		1 0.0331
LUNGTUT03	lung tumor, 69 M, match to LUNGNOT15		1 0.0310
BLADTUT02	bladder tumor, 80 F, match to BLADNOT03		1 0.0308
BRAITUT08	brain tumor, astrocytoma, 47 M		1 0.0305
PROSTUT12	prostate tumor, 65 M, match to PROSNOT20		2 0.0293
BLADNOT04	bladder, 28 M		1 0.0279 1 0.0278
TESTTUT02	testicular tumor, 31 M		
THYRNOT03	thyroid tumor, adenoma, 28 F		1 0.0278
SINTNOT13	small intestine, ileum ulcerativo chelitic as n		2.0.0277
COLNTUT03	colon tumor, 62 M, match to COLNNOT16		1 0.0275
BLADTUT05	bladder tumor, 66 M, match to BLADNOT06		1 0.0272
KIDNTUT01	kidney tumor, Wilms, 8m F		1 0.0268
PENITUT01	penis tumor, carcinoma, 64 M		1 0.0267 1 0.0267
COLNNOT23	colon, 16 M		1 0.0267
BRAITUT13	brain tumor, meningioma, 68 M		
LIVRTUT01	liver tumor, metastasis, 51 F		
PROSNOT14	prostate, 60 M, match to PROSTUTO8		1 0.0259 1 0.0256
BRSTTUT08	breast tumor, 45 F, match to BRSTNOT09		1 0.0254
BMARNOT03	bone marrow, 16 M		1 0.0234
RATRNOT02	heart, right atrium, 39 M		1 0.0236
PANCNOT01	pancreas, 29 M		1 0.0236
LUNGNOT04	lung, 2 M		0.0214
SYNORAT04	synovium, wrist, rheumatoid, 62 F		0.0174
PLACNOT02	placenta, fetal F		0.0168
BRSTNOT03	breast, 54 F, match to BRSTTUT02		L 0.0147
SPLNNOT04	spleen, 2 M		0.0128
PROSNOTO6	prostate, 57 M, match to PROSTUT04		0.0128
LUNGFET03	lung, fetal F	-	0.00114

Electronic Northern Results returned a total of 36 row(s).

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  1
                 М
                MPKGNKKPNEKKEELEKFAKELQGSDSDED SEQ ID NO-6
               G L L E T E P L Q G T D - E D A V A S A D F S S M L S E E E SEQ ID NO-1
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A V V I E Q P T V E P K L P Q N D S S S S N K I V L S Q A E SEQ ID NO-6
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              KEELKAELVQLEDEITTLRQVLSAKERHLV SEQ ID NO-1
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  36
              EIKQKLGMNLMNELKQNFSKSWHDMQTTTA SEQ ID NO-1
EIKRKLGINSLQELKQNIAKGWQDVTATSA SEQ ID NO-5
ELKRKLGLTPFSELSODINRSLKTVTDTDA SEQ ID NO-6
  66
 96 YKKTHETLSHAGQKATAAF - - - - SNVGTA SEQ ID NO-1
96 YKKTSETLSQAGOKASAAF - - - - SSVGSV SEQ ID NO-5
121 CTHFIEINIOKKKKQSMYYIKRLSKNIQTV SEQ ID NO-6
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- - I T K K L E D - - - - - SEQ ID NO-5
 151 PILTSEKKRILHAFIVLKKKSSILKSLLLW SEQ ID NO-6
 148 K S F E E R V E T T V T S L K T K V G G T N P N G G S F E E SEQ ID NO-1
135 KSFEEKVE---NLKSKVGGTKPAGGDFGE SEQ ID NO-5
211 KSFESKLGSALNN--AKMAASTSI----D SEQ ID NO-6
178 V L SSTAHASAOS L A G G S R R T K E E E L Q C
161 V L N S A A N A S A T T T E P L P E K T - Q E S L
234 H L A G A A R G P S O T G T P V A E E A K P I S
                                                                                                                                                                                                                         SEQ ID NO-1
                                                                                                                                                                                                                         SEQ ID NO-5
                                                                                                                                                                                                                         SEQ ID NO-6
```

FIGURE 5

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MTLFHFGNCFALAYFPYFITYKCSGLSEYN SEQ ID NO-3

MSFFHFTNCFALAFAPYFIVYKYSGINEYS SEQ ID NO-7

AFWKCVQAGVTYLFVQLCKMLFLATFFPTW SEQ ID NO-3

SIWKCATASGGYLLTOLAKLLITATFFPAL SEQ ID NO-7

EGGIYDFIGEFMKASVDVADLIGLNLVMSR SEQ ID NO-7

DSEGFSIVPEFLKSSADIIDVIGLHLLMTN SEQ ID NO-7

- NAGKGEYKIMVAALGWATAELIMSRCIPL SEQ ID NO-7

PLAGKGEVRFVVGGLGWGFAHSVAHRLVLL SEQ ID NO-7

120 WVGARGIEFDWKYIQMSIDSNISLVHYIVA SEQ ID NO-7

121 WVGARGTAFTWRWVOTSLDSSADLLVIVSL SEQ ID NO-7

150 SAQVWMITRYDLYHNFRPAVLLLMFLSVYK SEQ ID NO-7

150 SAQVWMITRYDLYHNFRPAVLLLMFLSVYK SEQ ID NO-7

180 AFVMETFVHLCSLGSWARLDAR--AVVTG SEQ ID NO-7

180 AFVMETFVHLCSLGSWARLDAR--SVYTG SEQ ID NO-7

180 AFVMETFVHLCSLGSWARLDAR--SVYTG SEQ ID NO-7

207 LLALKHFGPVCRRCQCALLGLVSOTLMYLF SEQ ID NO-7

218 PASLOVLVK SEQ ID NO-7
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FIGURE 6

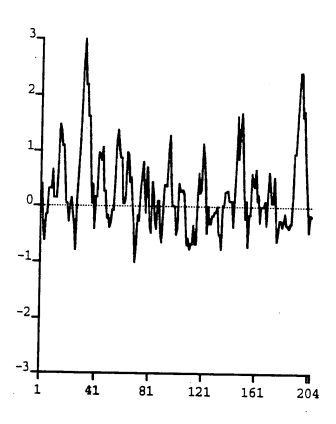


FIGURE 7

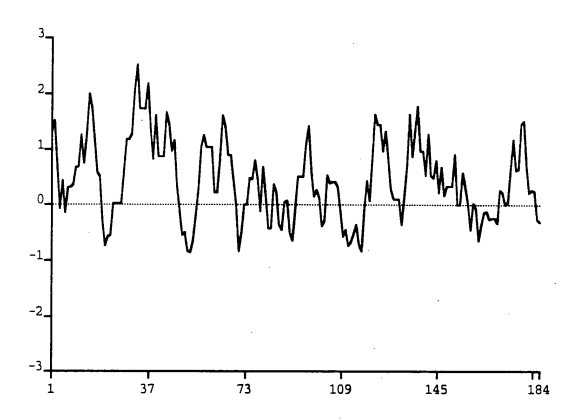


FIGURE 8

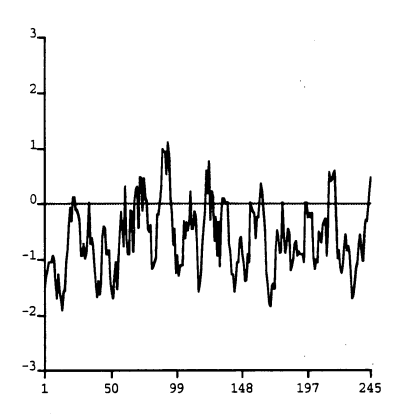


FIGURE 9

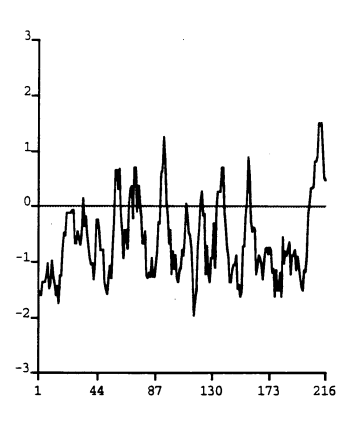


FIGURE 10

6043343

54	CAG	
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	AAG	
45	999	
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	GCT	
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GAA AGG CAT CTA GTT GAG ATA AAA CAA AAA CTC GGC ATG AAC CTG ATG AAT GAA	æ			TTA AAA CAG AAC TTC AGC	X
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APPROVED	O.G. FIG.			
. YB	CLASS	SUBCLASS		
DRAFTSMAN				

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FIGURE 1B

DOLLESSY . DORSOS

DRAFTSMAIL

774 783 NTG CCT GTG CTT ATC CAG ATA AGA AGA CCA AA 3'

APPROVED O.G. FIG.

BY CLASS SUBCLASS

DRAFTSMAN

AAA gcg GTG45 GCC CTG CCC CAG $^{\mathrm{TGC}}$ ලලල GGT CAG CGC င္လင္ပင 255 525 MKC

108 CGC CCG CGA 99 TCGသည gcc 90 GAC GGA CCC 81 GCT သည GGA GGA GAG ggg 63 CCG

162 TTC TGC AAC z GGG153 TTC CAC TTT144 CTG 口 ACC ATG ATC ggc ပ္သပ္ပ 126 CGG GAC 117 CGG GGG

216 GAG E 1 CC CTG GGC 207 ဗ AGC വ $_{\mathrm{TGC}}$ AAG 198 TAC ACC ATC 189 TTCTAC သည 180 CCC

270 CAA GTC TTTCTC261 TAC ACC GIC 252 GGA ტ GCTØ CAG GTC TGC AAA K 234 g_{CC} 225 AAC

324 ATC I ggc ggCGAA 315 306 TTTTTC ပ္သင္ဟ TTG 288 TTC F CTG AAG

378 ATA GAC 369 GTG GAT 360 AGC ATG M TTC GAG 342 999 TTC 333 GAC

FIGURE 2A

APPROVED	0.G. FIG.				
3Y	CLASS	SUBCLASS			
DRAFTSMAH	anin sakajuri made des 10 de das				

		•				
432 ATG M	486 CCC P	540 AGC S	594 TGG W	648 CTG L	702 TGC C	756 CTG L
ATC	ATT I	ATG M	GTC V	CTG	CTC	CTG
AAG K	TGC	CAG Q	CAG Q	CTT	CAC H	999 9
423 TAC Y	477 CGC R	531 ATC I	585 GCT A	639 GTC V	693 GTC V	747 ACG T
GAG	TCC	TAC Y	TCT	GCT	TTC	GTA V
GGA G	ATG M	AAG K	GCG A	CCA P	ACC	GTG V
414 AAG K	468 ATT I	522 TGG W	576 GTC V	630 CGG R	684 GAG E	738 GCA A
9	CTT	GAC D	ATC	TTC	ATG	CGA
GCC	GAG E	TTT F	TAC	AAC N	GTT V	GCC A
405 AAT N	459 GCT A	513 GAG E	567 CAT H	621 CAC H	675 TTT F	729 MTG X
CGG R	ACT	ATT I	GTC V	TAC Y	GCC	CTA
TCC	GCC	299	CTG L	CTG L	AAG K	RCT X
396 ATG	450 TGG W	504 CGG R	558 AGT S	612 GAT D	666 TAC Y	720 GCA A
GTC	၁၅၅	GCC	ATC	TAT Y	GTC	TGG W
CTT	CTG	GGA	AAC	CGC	AGT S	AGT
387 AAC N	441 GCC A	495 GTC V	549 TCC S	603 ACA T	657 CTC L	711 GGC G
CTA	GCT A	TGG W	GAC	ATA I	TTC	CTG
GGT CTA	GTT V	CTA TGG L W	ATA I	ATG M	ATG	TCG

FIGURE 2B

deleser loges

810 GGC

CTY CTA

765 AAG

CTC

ည္ဟ

TTC837 CTT

TAC ATG

864 GTG V

TTA

855 CAG (

CTC

TCA

CCT

846 GCC

GTT V

828 TTG

CAG ACA

Q

819 TCT S

GTG

TTG

GGA AAG 882 ACA GTA TTT 873 TAA

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AAG K

APPROVED	0.G. F	IG.		
BY:	CLASS	SUBCLASS		
DRAFTSMAH				

Library	Lib Description	Abun	Pct Abun
1 1 1 1 1 1 1 1		1 1 1 1 1 1 1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
BRAINOM02	brain, 55 M, NORM, WM	H	0.0454
UTRSNOT01	uterus, 59 F	\leftarrow	0.0393
TLYMNOR01	lymphocytes (non-adher PBMNC), 24 M, RP	· ~	0.0372
BRSTNOT02	breast, 55 F, match to BRSTTUT01	2	0.0317
PROSTUT08	prostate tumor, 60 M, match to PROSNOT14	-	0.0266
PROSNOT14	prostate, 60 M, match to PROSTUT08	⊣	0.0256
LIVRNOM01	liver, 49 M, WM	~	0.0254
PROSNOT15	prostate, 66 M, match to PROSTUT10	\leftarrow	0.0241
NERVMSM01	multiple sclerosis, 46 M, NORM, WM	⊣	0.0228
HNT2AGT01	hNT2 cell line, post-mitotic neurons	⊣	0.0190
BRAITUT02	brain tumor, metastasis, 58 M	Н	0.0169
LIVSFEM02	liver/spleen, fetal M, NORM, WM	7	0.0053

Electronic Northern Results returned a total of 12 row(s).

FIGURE 3

Library	Lib Description	Abun	Pct Abun
COLNNOT22	colon, 56 F	5	0.0554
COLNPOT01	colon polyp, 40 F	7	0.0513
PROSNOT18	prostate, hyperplasia, 58 M	2	0.0513
MUSCNOT02	muscle, psoas, 12 M	Н	0.0382
STOMTUT01	stomach tumor, 52 M, match to STOMNOT02	1	0.0368
SINTNOT02	small intestine, 55 F	٦	0.0337
LVENNOT03	heart, left ventricle, 31 M	Н	0.0337
MMLR3DT01	macrophages (adher PBMNC), M/F, 72-hr MLR	Н	0.0331
PROSTUT01	prostate tumor, 50 M, match to PROSNOT02	Н	0.0310
LUNGTUT03	lung tumor, 69 M, match to LUNGNOT15	7	0.0308
BLADTUT02	bladder tumor, 80 F, match to BLADNOT03	7	0.0305
BRAITUT08	brain tumor, astrocytoma, 47 M	7	0.0293
PROSTUT12	prostate tumor, 65 M, match to PROSNOT20	Н	0.0279
BLADNOT04	bladder, 28 M	П	0.0278
TESTTUT02	testicular tumor, 31 M	Н	0.0278
THYRNOT03	thyroid tumor, adenoma, 28 F	7	0.0277
SINTNOT13	small intestine, ileum, ulcerative cholitis, 25 F	⊣	0.0275
COLNTUT03	colon tumor, 62 M, match to COLNNOT16	⊣	0.0272
BLADTUT05	bladder tumor, 66 M, match to BLADNOT06	⊣	0.0268
KIDNTUT01	kidney tumor, Wilms, 8m F	Н	0.0267
PENITUT01	penis tumor, carcinoma, 64 M	Н	0.0267
COLNNOT23	colon, 16 M	Н	0.0264
BRAITUT13	brain tumor, meningioma, 68 M	Н	0.0262

FIGURE 4A

APPROVED	O.G. FIG.			
; BY	CLASS	SUBCLASS		
DRAFTSMAR				

LIVRTUTUL	liver tumor, metastasis, 51 F	.	0.0259	. •
PROSNOT14	prostate, 60 M, match to PROSTUT08	Н	0.0256	
BRSTTUT08	breast tumor, 45 F, match to BRSTNOT09	Н	0.0254	
BMARNOT03	bone marrow, 16 M	H	0.0242	
RATRNOT02	héart, right átrium, 39 M	\leftarrow	0.0236	
PANCNOT01	pancreas, 29 M	₽	0.0214	
LUNGNOT04	lung, 2 M	⊢	0.0183	
SYNORAT04	synovium, wrist, rheumatoid, 62 F	· ႕	0.0174	
PLACNOT02	placenta, fetal F		0.0168	
BRSTNOT03	breast, 54 F, match to BRSTTUT02	П	0.0147	
SPLNNOT04	spleen, 2 M	~	0.0128	
PROSNOT06	prostate, 57 M, match to PROSTUT04	Т	0.0114	
LUNGFET03	lung, fetal F	⊣	0.0091	

Electronic Northern Results returned a total of 36 row(s)

FIGURE 4B

	- EDAVASADFSSMLSEEEKEELKAELVQLEDEITTLRQ SEQ ID NO-1 - EDVAATISATETLSEEERRELAKVEEEIQTLSQ SEQ ID NO-5 LPQNDSSSNKIVLSQAEKDLLRTELDKTEEISTLKQ SEQ ID NO-6	SAKERHLVEIKQKLGMNLMNELKQNFSKSWHDMQTTTA SEQ ID NO-1 AAKEKHLAEIKRKLGINSLQELKQNIAKGWQDVTATSA SEQ ID NO-5 SARQKHAAELKRKLGLTPFSELSQDINRSLKTVTDTDA SEQ ID NO-6	K T K T K T K T K T K T K T K T K T K T	 A F I V	-MRNSPTFKSFEERVETTVTSLKTKVGGTNPNGGSFEE SEQ ID NO-1 -VKNSPTFKSFEEKVE NLKSKVGGTKPAGGDFGE SEQ ID NO-5 DMRNSSLFKSFESKLGSALNN AKMAASTSI D SEQ ID NO-6	SSTAHASAOSLAGGSRRTKEEELQC NSAANASATTTEPLPEKT-QESL
M M W W W W W W W W W W W	TD-EDA EG-EDV PKLPQN	저 저 저	YKKTHE YKKTSE CTHFIE		N N N	N N C

FIGURE :

\vdash	MTLFHFGNCFALAYFPYFITYKCSGLSEYNAFWKCVQAGV SEQ ID NO-3 MSFFHFINCFALAFAPYFIVYKYSGINEYSSIWKCATASG SEQ ID NO-7
44	TYLFVQLCKMLFLATFFPTWEGGIYDFIGEFMKASVDVAD SEQID NO-3 GYLLTOLAKLLIIATFFPALDSEGFSIVPEFLKSSADIID SEQID NO-7
ਲ ਲ	LIGLNLVMSR-NAGKGEYKIMVAALGWATAELIMSRCIPL SEQID NO-3 VIGLHLLMTNFLAGKGEVRFVVGGLGWGFAHSVAHRLVLL SEQID NO-7
120	O WVGARGIEFDWKYIQMSIDSNISLVHYIVASAQVWMITRY SEQID NO-3 1 WVGARGTAFTWRWVOTSLDSSADLLVIVSLACLTWMITR - SEQID NO-7
160 160	160 DLYHNFRPAVLLLMFLSVYKAFVMETFVHLCSLGSWARLD SEQID NO-3 160TPNKFLVSPILAITVOHTFSLYGWSLLA SEQID NO-7
200	200 AR AVVTGLLALKHFGPVCRRCQCALLGLVSOTLMYLF SEQID ND-3 188 FRFAYSIATAILTVVVYSA NRTASTRKN SEQID ND-7

SEQ ID NO-3 SEQ ID NO-7

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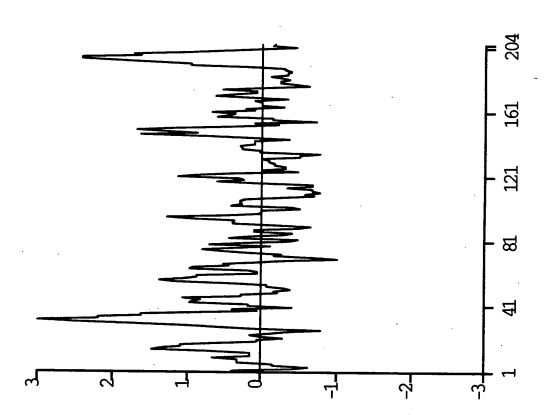
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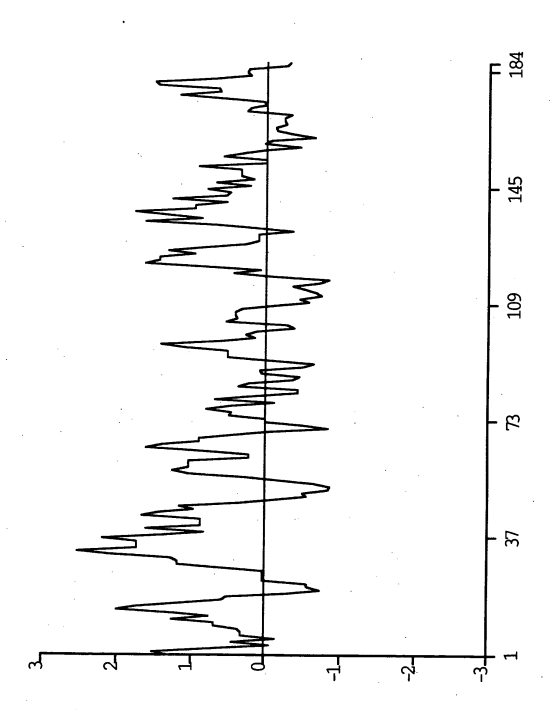


FIGURE 8

APPROVED	O.G. FIG.		
BY	CLASS	SUBCLASS	
DRAFTSHAN			

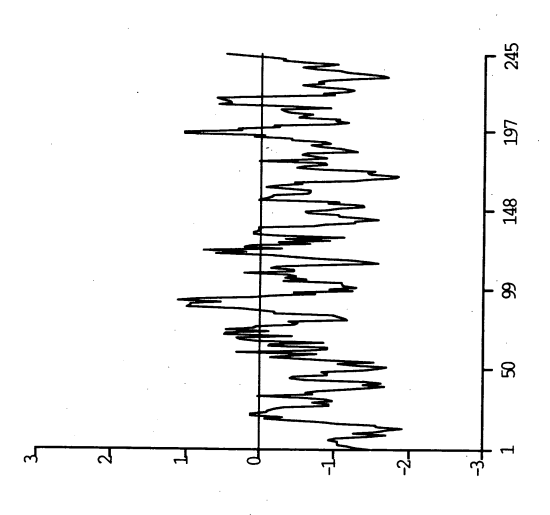
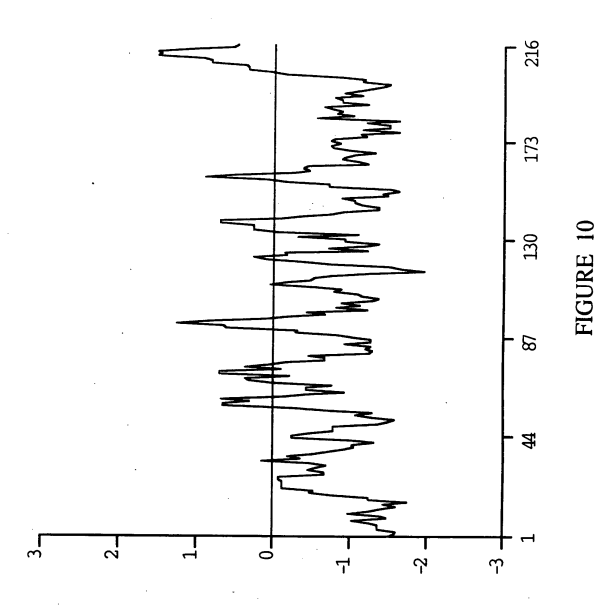


FIGURE 9





aeca. Zezate